



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁵ : A61K 6/05, 6/00	A1	(11) International Publication Number: WO 92/11837 (43) International Publication Date: 23 July 1992 (23.07.92)
(21) International Application Number: PCT/US91/08809 (22) International Filing Date: 25 November 1991 (25.11.91) (30) Priority data: 638,614 8 January 1991 (08.01.91) US (71) Applicant: MINNESOTA MINING AND MANUFACTURING COMPANY [US/US]; 3M Center, Post Office Box 33427, Saint Paul, MN 55133-3427 (US). (72) Inventors: OXMAN, Joel, David ; FUNDINGSLAND, Jon, Wallace ; Post Office Box 33427, Saint Paul, MN 55133-3427 (US). (74) Agents: JOSEPHS, Barry, D. et al.; Minnesota Mining and Manufacturing Company, Post Office Box 33427, Saint Paul, MN 55133-3427 (US).		(81) Designated States: AT (European patent), AU, BE (European patent), BR, CA, CH (European patent), DE (European patent), DK (European patent), ES (European patent), FR (European patent), GB (European patent), GR (European patent), IT (European patent), JP, KP, LU (European patent), NL (European patent), PL, SE (European patent), SU . Published <i>With international search report.</i>
(54) Title: ADHESIVE AMALGAM SYSTEM (57) Abstract A modified amalgam composition forming an adhesive bond with tooth structure treated with a dental adhesive. The modified amalgam can be prepared by admixing particulate additives into conventional amalgam alloy powder to form a modified alloy powder and then triturating the modified alloy powder with mercury. The modified amalgam when applied to a prepared tooth cavity that has been precoated with an acrylate- or methacrylate-functional dental adhesive results in an adhesive bond between the modified amalgam and coated tooth structure. Preferred particulate additives for the amalgam alloy powder are acrylate- or methacrylate-functional polymers, metal salts of acrylates or methacrylates, nonmetallic fillers, oxidizing agents and reducing agents.		

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	ES	Spain	MG	Madagascar
AU	Australia	FI	Finland	ML	Mali
BB	Barbados	FR	France	MN	Mongolia
BE	Belgium	GA	Gabon	MR	Mauritania
BF	Burkina Faso	GB	United Kingdom	MW	Malawi
BG	Bulgaria	GN	Guinea	NL	Netherlands
BJ	Benin	GR	Greece	NO	Norway
BR	Brazil	HU	Hungary	PL	Poland
CA	Canada	IT	Italy	RO	Romania
CF	Central African Republic	JP	Japan	RU	Russian Federation
CG	Congo	KP	Democratic People's Republic of Korea	SD	Sudan
CH	Switzerland	KR	Republic of Korea	SE	Sweden
CI	Côte d'Ivoire	LI	Liechtenstein	SN	Senegal
CM	Cameroon	LK	Sri Lanka	SU	Soviet Union
CS	Czechoslovakia	LU	Luxembourg	TD	Chad
DE	Germany	MC	Monaco	TG	Togo
DK	Denmark			US	United States of America

ADHESIVE AMALGAM SYSTEMBACKGROUND OF THE INVENTION

5

1. Field of the Invention

The present invention relates to amalgam compositions for restorative dental repair.

10 2. Description of the Prior Art

Dental amalgam has been available to the dental profession for well over a century and it is used extensively for intracoronal and extracoronal restorations. Amalgam is highly durable and the strength and occlusal wear characteristics of alternative materials such as composite resins are generally compared to that of amalgam. However, amalgam does not adhere to tooth structure and the dentist must take great care to prepare the tooth cavity with dovetails and various cutout grooves which in effect mechanically lock the amalgam into the cavity. Such required preparation by the dentist results in the need to excavate more tooth structure than would otherwise be necessary if the amalgam were adhesive. This of course weakens the tooth. Additionally the problem of microleakage at the interface of the amalgam and cavity wall tends to occur for a period after the amalgam has been placed into the cavity. Microleakage allows penetration of bacteria, soluble salts and saliva into any space between the amalgam restoration and cavity walls. This can lead to inflammation and pulp irritation which in turn can cause other complications. The penetration of bacteria into spaces between the amalgam and cavity wall can demineralize the cavity walls and lead to formation of recurrent caries. Corrosion of amalgam can cause one of the amalgam alloy metals, for example tin, to deposit along the amalgam and cavity wall interface. This

- 2 -

usually leads to tooth discoloration along the interface and can noticeably detract from the esthetic appearance of tooth and restoration. An adhesive seal between amalgam restoration and cavity walls could prevent
5 microleakage. An adhesive amalgam could significantly reduce the amount of tooth the dentist needs to excavate in order to prepare the cavity for restoration. An adhesive amalgam could impart significantly more strength to the filled tooth structure.

10 The concept of making an adhesive amalgam is thus attractive from many standpoints. Unfortunately the nature of amalgam and teeth makes it very difficult to adhere amalgam to tooth structure.

U.S. Pat. No. 3,513,123 (Saffir) discloses an
15 epoxy liquid resin composition which is added to amalgam in an effort to make the amalgam adhere to tooth structure. This reference discloses use of an epoxy liquid resin additive consisting of a glycidyl ether type epoxy resin containing a polyamine hardening agent.

20 Various references disclose mixtures of amalgam with various additives to impart improved mechanical characteristics. For example U.S. Pat. No. 4,859,412 (Grell) discloses the addition of ceramic or glass powder to alloy powder, which when amalgamated with mercury
25 produces an amalgam with improved mechanical strength properties. Japanese patent publication 55-22545 discloses use of glass additives which can be blended with amalgam during the amalgamation of alloy powder with mercury. The amalgam modified with glass additives is
30 alleged to impart improved compressive strength properties.

U.S. Pat. No. 2,991,176 (Clancy) discloses mixtures of silica powder, alumina and other materials with amalgam alloy powder. The amalgam alloy powder and
35 other materials are milled to form particles wherein these materials are encapsulated by the amalgam alloy powder. When the encapsulated particles are amalgamated

- 3 -

with mercury, a modified amalgam is formed and is said to have reduced thermal coefficient of expansion which in turn reduces the chance for microleakage.

U.S. Pat. Nos. 4,255,192 (Burns), 4,684,347
5 (Palaghias) and 3,676,112 (Muhler) disclose modified amalgams which utilize various additives or treatment of the amalgam alloy powder to impart varying physical or cariostatic properties to the amalgam.

U.S. Pat. No. 4,064,629 (Stoner), discloses a
10 method for applying amalgam restorations which involves precoating the surfaces of a cavity within a carious tooth with a layer of an "adhesive-metal" lining composition. The metal of the lining composition is amalgamated by diffusion of the mercury from the
15 subsequently applied conventional dental amalgam filling. The "adhesive-metal" lining composition is said to improve corrosion resistance of the dental amalgam filling and also promotes bonding between the amalgam restoration and the cavity surfaces. Other references
20 which disclose precoating the surfaces of a tooth cavity with an adhesive coating said to adhere to conventional amalgam are, for example, U.S. Pat. Nos. 4,001,483 (Lee) and 3,574,943 (Nicholson).

In recent years several adhesive products which
25 claim to make amalgam adhesive to tooth structure have been made available to dental clinicians. (The term "tooth structure" as used hereinafter shall be interpreted to include either or both dentin and enamel, optionally precoated with liner or base.) One such
30 product is sold in a kit form under the trademark "AMALGAMBOND" available from Parkell Co. The "AMALGAMBOND" product is a liquid adhesive resin which is coated directly onto tooth structure. The application and curing procedure are cumbersome and involve a number
25 of steps. The curing procedure also requires use of an air-sensitive catalyst which if dropped on flammable paper causes smoldering. The active ingredients in the

- 4 -

adhesive are 4-META (4-methacryloxyethyl trimellitic anhydride) and TBB (tri-n butyl borane). Other products which similarly involve coating a specific curable resin directly onto tooth structure to make amalgam adhere are available under the trademarks "PANAVIA" Dental Adhesive from Kuraray Company and "SUPERBOND" Adhesive from Sun Medical Co., Ltd., Kyoto, Japan. These latter products also are difficult to employ, since there are a number of required preparatory steps for their application and curing.

Literature articles which disclose bonding of amalgam to tooth structure by precoating the tooth with adhesive resin include M. Staninec and M. Holt, Journal of Prosthetic Dentistry (1988), Vol. 59, p. 397-402, A. Lacey and M. Staninec, Quintessence International (1989), Vol. 20, p. 521-524, and Y. Torii, et al. Operative Dentistry (1989), Vol. 14, p. 142-148. The above listed articles report improved adhesive tensile strength between amalgam and coated tooth structure but do not report adhesive shear bond strength of the amalgam.

SUMMARY OF THE INVENTION

The present invention is directed in principal aspect to a modified amalgam composition which produces an adhesive bond with treated tooth structure. In the preferred embodiments the tooth structure is coated with an acrylate- or methacrylate-functional dental adhesive.

The modified amalgam of the invention is produced by admixing particulate additives into conventional amalgam. A modified amalgam of the invention can be produced, for example, by admixing particulate additives into "DISPERSALLOY" alloy powder. The modified amalgam is then prepared in a conventional manner by triturating the modified alloy powder with mercury in an amalgamator. The modified amalgam when applied to a prepared tooth cavity which has been

- 5 -

precoated with acrylate- or methacrylate-functional dental adhesive results in an adhesive bond between the modified amalgam and the coated tooth structure. Since the modified amalgam adheres to the coated tooth structure, the dentist generally will not need to excavate as much tooth structure in preparing the cavity as would be necessary when placing a conventional unmodified amalgam restoration. This results in a saving of tooth structure and reduces the chance of the tooth weakening because of the cavity preparation. Also, the present invention significantly reduces the chance of microleakage occurring at the interface between the amalgam and cavity walls, since the adhesive bond between amalgam and tooth structure discourages penetration of bacteria, soluble salts and saliva between the amalgam restoration and cavity walls.

The preferred particulate additives which can be admixed into conventional amalgam to produce the modified amalgam of the invention are selected from the following groups: 1) acrylate- or methacrylate-functional polymers, 2) metal salts of acrylates or methacrylates, 3) nonmetallic fillers, 4) oxidizing agents, and 5) reducing agents.

25 DETAILED DESCRIPTION

A preferred embodiment of the present invention is a restorative system for making amalgam adherent to tooth structure. The system of the invention involves production of a modified formulation for amalgam and employing that modified formulation in combination with dental adhesive applied to tooth structure. The modified amalgam composition of the invention involves adding a particulate material, preferably nonmetallic particulate material in powder form, to conventional dental amalgam alloy powder. As is well known, conventional amalgam preparations are available in capsules which contain

- 6 -

amalgam alloy powder and mercury, sealed by a penetrable bladder located at one end of the capsule. A small metal rod is included in the capsule. The clinician prepares amalgam by placing the capsule containing the alloy powder and mercury into an amalgamator. The amalgamator, (or triturator as it is often called), vibrates at high speed so that the metal rod within the capsule can penetrate the bladder to release the mercury contained therein. As mercury admixes into the alloy powder a reaction occurs between alloy powder and mercury and the amalgam slowly begins to set. At this stage the amalgam is ready for packing into the tooth cavity.

The present invention involves the simple addition of a particulate material to conventional amalgam alloy powder. The powder additives of the present invention are intended to be applicable to the full range of conventional amalgam alloy powders and conventional weight ratios of mercury in relation to total amalgam alloy powder. Conventional alloy powders are typically mixtures of silver, tin, copper, and zinc. Conventional amalgam alloy powders have proper proportioning of these metals to result in an alloy described in the art as a "balanced alloy." For example it is known that increasing the silver content increases the expansion of the setting amalgam, shortens setting time, increases compressive strength, and tends to make the alloy mixture more difficult to amalgamate. Tin behaves in an opposite way for all these properties. Copper and zinc contribute properties similar to silver with respect to expansion, setting time and strength, but copper is used principally for increased strength and zinc for increased resistance to tarnish. Conventional alloys are broadly classified as low-copper alloys (5% or less copper) and high copper alloys (13% to 30% copper). Commercially available low copper amalgam alloys contain typically the following compositions which apply to lathe-cut or spherical particle shapes: silver (63-70%),

- 7 -

tin (26-28%), copper (2-5%), and zinc (0-2%).

Commercially available high copper alloys using lathe-cut particles contain typically the following compositions:

silver (40-70%), tin (26-30%), copper (2-5%), and zinc

5 (0-2%). Commercially available high copper alloys using spherical particles contain typically the following compositions: silver (40-65%), tin (0-30%), copper (20-40%), zinc (0%), and palladium (0-1%). Mercury typically represents 40 to 60 percent by weight of the amalgam mix.

10 A widely used high copper amalgam alloy is available under the trademark "DISPERSALLOY" alloy from Johnson & Johnson Company. The "DISPERSALLOY" alloy contains about 13% copper and the mixed "DISPERSALLOY" amalgam contains about 50% mercury.

15 We have discovered that the particulate additives of the invention when mixed into conventional amalgam alloy powder (preferably high copper alloy powder, e.g. "DISPERSALLOY" alloy powder) provide a modified restorative amalgam that adheres very

20 effectively to tooth structure which has been precoated with a dental adhesive. Representative preferred dental adhesives for the system of the invention are available under the trademark "SCOTCHBOND" Dual Cure Dental Adhesive and "SCOTCHBOND 2" Light Cure Dental Adhesive,

25 both from 3M Dental Products Division. "SCOTCHBOND" Dual Cure Dental Adhesive is described in U.S. Pat. Nos. 4,544,467 (Bunker), 4,670,576 (Bunker), 4,929,746 (Bunker), and 4,669,983 (Bunker). The compounds disclosed in these references include generally acrylate-

30 or methacrylate-functional dental adhesives which bond to tooth structure. The "SCOTCHBOND" Dual Cure Dental Adhesive can be applied to tooth structure and cured with light or under room temperature conditions in the manner described in the "SCOTCHBOND" Dual Cure Dental Adhesive

35 kit.

The composition of the "SCOTCHBOND 2" Light Cure Dental Adhesive is disclosed in U.S. Pat. No.

- 8 -

4,719,149 (Aasen). The "SCOTCHBOND 2" dental adhesive is applied to tooth structure and light cured in the manner described in the "SCOTCHBOND 2" Light Cure Dental Adhesive kit. Both "SCOTCHBOND" and "SCOTCHBOND 2" dental adhesives were commonly intended for bonding various conventional composite restorative resins to tooth structure.

Other preferred dental adhesives which can be employed with the modified amalgam of the present invention contain acrylate- or methacrylate-functional polymers and may also contain phosphorous compounds. In such dental adhesives either a single phosphorus compound or a mixture of phosphorus compounds can be used. Dental adhesives which can be precoated onto tooth structure and used with the modified restorative amalgam of the invention to produce an adhesive bond between the amalgam and tooth structure include "ALL-BOND" Universal Dental Adhesive System from Bisco, Inc., "CLEARFIL" Photo Bond Light Cured Dental Bonding Agent from Kuraray Co., Ltd., "RESTOBOND 3" Dual Dentin-Enamel Bonding Agent from Lee Pharmaceuticals, (see U.S. Pat. Nos. 4,524,527 and 4,521,550), "TENURE" Solution Dentin Bonding System from Den-Mat Corporation, "GLUMA" Bonding System from Columbus Dental Miles, Inc., "PRISMA UNIVERSAL BOND 2" Dentin/Enamel Bonding Agent from L. D. Caulk Division of Dentsply International, Inc., (see U.S. Pat. No. 4,814,423), "MIRAGE-BOND" Dentin-Enamel Bonding System from Chameleon Dental Products, Inc., (see U.S. Pat. Nos. 4,514,527, 4,521,550, 4,588,756, and 5,659,751), "BONDLITE" dental adhesive from Sybron Corp., "Johnson & Johnson" dentin bonding agent and "Johnson & Johnson" light-curing bonding agent, both from Johnson & Johnson Co., "PALFIQUE" bonding agent from Tokuyama Soda Co., Ltd., "SHOFU" bonding base from Shofu, Inc, and "SINTERBOND" dental adhesive from Teledyne Getz. All of the above adhesives are acrylate- or methacrylate-functional dental adhesives.

- 9 -

If desired, other free-radically polymerizable non-phosphorus-containing compounds can be mixed with the dental adhesive, for example, as a diluent to reduce viscosity or promote wetting. Other suitable free-radically polymerizable compounds include mono- or poly- (e.g., di-, tri- or tetra-functional) acrylates and methacrylates such as methyl acrylate, 2-hydroxyethyl acrylate, triethyleneglycol diacrylate, neopentylglycol diacrylate, hexamethyleneglycol diacrylate, trimethylolpropane triacrylate, pentaerythritol tetraacrylate, polyalkylene glycol mono- and diacrylates, urethane mono- or poly-functional acrylates, Bisphenol A diacrylates, and the corresponding methacrylates of the above compounds, as well as acrylamides and methacrylamides, vinyl compounds, styrene compounds, and other olefinically unsaturated compounds suitable for use in the oral environment. U.S. Pat. Nos. 4,499,251, 4,515,930, 4,537,940 and 4,539,382 contain an extensive list of such compounds.

We have found that the following groups of additives when added to conventional amalgam alloy powder, e.g., "DISPERSALLOY" powder, make the amalgam strongly adherent to tooth structure precoated with an acrylate- or methacrylate-functional dental adhesive.

Group 1 - Acrylate- or Methacrylate-Functional Polymers

Representative acrylate- or methacrylate-functional moieties include poly(alkanoic acid) powder. This polymer is a copolymer of itaconic and acrylic acid, and is described in European published patent application No. 88-312127.

Group 2 - Metal Salts of Acrylates or Methacrylates

Representative salts from this group include zinc dimethacrylate, zirconium dimethacrylate, silver methacrylate, sodium methacrylate, and magnesium methacrylate.

Group 3 - Nonmetallic Fillers

- 10 -

Nonmetallic fillers include both untreated organic fillers and surface-treated fillers.

Representative nonmetallic fillers include blends of 66 wt% "OX-50" pyrogenic silica available from Degussa

5 Company, 17 wt% tetraethyleneglycol dimethacrylate ("TEGDMA") from Rohm Tech Co. and 17 wt% diglycidylether dimethacrylate ("BIS-GMA"). (The "OX-50" pyrogenic silica was treated with 20 wt% gamma methacryloxypropyl trimethoxysilane.) The blend is thermally polymerized to

10 a hard mass which is reground to yield a fine powdered filler with particle size less than 50 microns. (The surface of the "OX-50" pyrogenic silica particles could alternatively be treated with gamma-mercaptopropyltriethoxysilane or gamma-

15 aminopropyltrimethoxysilane). Other representative nonmetallic fillers include zirconia/silica filler pretreated with gamma-methacryloxypropyl trimethoxysilane as described in U.S. Pat. No. 4,503,169.

Group 4 - Oxidizing Agents

20 Preferred oxidizing agents include benzoyl peroxide.

Group 5 - Reducing Agents

Preferred reducing agents include sodium benzenesulfinate.

25 The above-listed additives either alone or in any combination are preferably admixed into the amalgam alloy powder. Alternatively, where convenient, the additives can be added to the mercury. These additives may also be admixed into the amalgam prepared from alloy
30 powder and mercury just after trituration.

Examples 1-17 show specific powder additives which we have used to modify the alloy. The amount of mercury used in the amalgam is also shown. The specific additive powder shown in Examples 1-17 was added to
35 "DISPERSALLOY" alloy powder which was contained in a conventional amalgam capsule. The capsule contained mercury protected by a penetrable bladder. The capsule

- 11 -

in each case was then closed and vibrated in an amalgamator which upon trituration formed the modified restorative amalgam.

The protocol for preparing the bovine teeth and measuring shear bond strength is as follows. Bovine teeth of similar age and appearance were partially embedded in circular acrylic discs. The exposed portion of each tooth was ground flat and parallel to the acrylic disc using Grade 120 silicon carbide paper-backed abrasive mounted on a lapidary wheel, in order to expose the enamel. During this and subsequent grinding and polishing steps, the teeth were continuously rinsed with water. Further grinding and polishing of the teeth was carried out by mounting Grade 600 silicon carbide paper-backed abrasive on the lapidary wheel. The polished teeth were stored in distilled water, and used for testing within 2 hours after polishing. The polished teeth were removed from the water and dried using a stream of compressed air.

Phosphoric acid etching gel was applied to the exposed enamel for 15 seconds, rinsed with water and dried. "SCOTCHBOND" Dual Cure Adhesive or "SCOTCHBOND 2" Dental Adhesive as indicated in the Examples, was then applied to the exposed enamel with a brush and blown into a thin film with compressed air and then cured with a "VISILUX" 2 dental curing light. The modified restorative amalgam was then packed onto the precoated bovine tooth structure. The adhesive shear bond strength of the modified amalgam was then measured as described below.

Previously prepared molds made from a 2 mm thick "Teflon" sheet with a 5 mm diameter hole through the sheet were clamped to each polished tooth so that the central axis of the hole in the mold was normal to the polished tooth surface. The hole in each mold was filled with a specific modified amalgam of formulation shown in Table I. The teeth and molds were allowed to stand for

- 12 -

about 15 minutes at room temperature, then stored in distilled water at 37°C. for 24 hours. The molds were then carefully removed from the teeth, leaving a molded button of amalgam attached to each tooth.

5 Adhesive strength was evaluated by mounting the acrylic disk in a holder clamped in the jaws of an "Instron" apparatus with the polished tooth surface oriented parallel to the direction of pull. A loop of orthodontic wire (0.44 mm diameter) was placed around the
10 restorative button adjacent to the polished tooth surface. The ends of the orthodontic wire were clamped in the pulling jaw of the Instron apparatus, thereby, placing the bond in shear stress. The bond was stressed until it or the button failed, using a crosshead speed of
15 2 mm/min.

Comparative Examples A-D were run using a modified amalgam, but without coating the bovine tooth with dental adhesive. In these Comparative Examples the modified amalgam was prepared by mixing various
20 particulate additives into conventional amalgam alloy powder, e.g. "DISPERSALLOY" alloy, and amalgamating the mixture for about 20 seconds. In each Comparative Example the adhesive shear bond strength of the modified amalgam was 0 kg/cm². The modified amalgam fell off the
25 tooth structure before placement in water.

Control A and B were prepared using unmodified amalgam applied to precoated tooth structure. In Control A amalgam prepared using unmodified "DISPERSALLOY" amalgam alloy powder was applied to bovine tooth
30 structure precoated with "SCOTCHBOND" Dual Cure Dental Adhesive. The average shear bond strength was 15 kg/cm². In Control B amalgam prepared using unmodified "DISPERSALLOY" amalgam alloy powder was applied to bovine tooth structure precoated with "SCOTCHBOND 2" Dental
35 Adhesive. The average shear bond strength was 0 kg/cm².

Table I gives a tabular listing of the adhesive shear bond strength of "DISPERSALLOY" amalgam alloy

- 13 -

powder modified by the addition of the respective powder additive in each Example given in weight percent of total amalgam. In all cases the amount of mercury present in the total amalgam mix equaled the amount of

- 5 "DISPERSALLOY" alloy powder present in the mix. The adhesive used to precoat the prepared bovine tooth structure in the manner as above described is also listed in Table I. In all cases except Examples 7, 8 and 14, the adhesive shear bond strength of the modified amalgam
10 of the invention was greater than that of the unmodified (Control A and B) "DISPERSALLOY" amalgam.

TABLE I
ADHESIVE SHEAR BOND STRENGTH OF
MODIFIED AMALGAM APPLIED TO PREPARED TOOTH STRUCTURE

EXAMPLE NUMBER	TOOTH STRUCTURE ADHESIVE PRECOAT	ADDITIVE TO "DISPERSALLOY" ALLOY POWDER	ADDITIVE IN AMALGAM ¹ (WT%)	AVERAGE ADHESIVE SHEAR BOND STRENGTH (kg/cm ²)
1	"SCOTCHBOND" Dual Cure Dental Adhesive	Poly(alkanoic acid) powder ²	2.00	26
2	"	Zinc dimethacrylate	0.25	25
3	"	"	0.50	28
4	"	"	1.00	66
5	"	"	2.00	50
6	"	Zirconium dimethacrylate	0.5	16
7	"	"	1.00	13
8	"	Silver methacrylate	0.38	13
9	"	Organic filler ³	0.25	55

-15-

TABLE I (cont.)

EXAMPLE NUMBER	TOOTH STRUCTURE ADHESIVE PRECOAT	ADDITIVE TO "DISPERSALLOY" ALLOY POWDER	ADDITIVE IN ANALGAM ¹ (WT%)	AVERAGE ADHESIVE SHEAR BOND STRENGTH (kg/cm ²)
10	"SCOTCHBOND" Dual Cure Dental Adhesive	Organic filler ³	0.50	72
11	"	"	1.00	25
12	"	"	2.00	24
13	"	Zirconia/silica filler ⁴	1.00	19
14	"	Zinc glass powder ⁵	2.00	7
15	"SCOTCHBOND 2" Dental Adhesive	Benzoyl peroxide	0.16	72
16	"SCOTCHBOND" Dual Cure Dental Adhesive	Benzoyl peroxide	0.38	43
17	"	Sodium benzenesulfinate	0.38	25

SUBSTITUTE SHEET

TABLE I (cont.)

EXAMPLE NUMBER	TOOTH STRUCTURE ADHESIVE PRECOAT	ADDITIVE TO "DISPERSALLOY" ALLOY POWDER	ADDITIVE IN AMALGAM ¹ (WT%)	AVERAGE ADHESIVE SHEAR BOND STRENGTH (kg/cm ²)
Comparative A	None	Zinc dimethacrylate	1.0	0
Comparative B	None	Organic filler ³	0.5	0
Comparative C	None	Zinc glass powder ⁵	2.0	0
Comparative D	None	Benzoyl peroxide	0.38	0
Control A	"SCOTCHBOND" Dual Cure Dental Adhesive	None	0	15
Control B	"SCOTCHBOND 2" Dental Adhesive	None	0	0

Notes:

1. Amalgam in all cases contained equal parts by weight mercury and "DISPERSALLOY" alloy powder (before addition of additives), e.g., composition of comparative Example B was "DISPERSALLOY" alloy powder (49.75 wt%), mercury (49.75 wt%) and organic filler (0.5 wt%).
2. poly(alkanoic acid) powder prepared according to Example 11 of European published patent application No. 88-312127.
3. Organic filler contained 66 wt% "OX-50" pyrogenic silica which has been treated with 20 wt% gamma-methacryloxypropyl trimethoxysilane, 17 wt% TEGDMA and 17 wt% BIS-GMA.
4. Zirconia/silica filler powder prepared according to Example 1 of U.S. Pat. No. 4,503,169.
5. Zinc glass powder prepared according to Example 13 of European published patent application No. 88-312127.

- 18 -

While the present invention has been described with respect to specific embodiments it should be appreciated that the invention is not intended to be limited to such embodiments. Chemical species, other than
5 the preferred species within a disclosed class of additives, may be substituted for the preferred without departing from the scope of the invention. Therefore, the present invention is not intended to be limited to the preferred embodiments but rather is defined by the claims
10 and equivalents thereof.

- 19 -

WE CLAIM:

1. A dental restorative system comprising a modified amalgam for application to tooth structure in combination with a dental adhesive for application to
5 tooth structure so that at least a portion of the modified amalgam shall contact tooth structure precoated with said dental adhesive, the modified amalgam comprising silver, mercury and particulate additive, said particulate additive having the property of rendering said modified
10 amalgam adherent to said precoated tooth structure.
2. A restorative system as in Claim 1 wherein the modified amalgam comprises an amalgamation of modified amalgam alloy powder and mercury, said modified amalgam
15 alloy powder comprising said silver and particulate additive.
3. A restorative system as in Claim 1 wherein the dental adhesive comprises an acrylate- or
20 methacrylate-functional resin.
4. A restorative system as in Claim 1 wherein the particulate additive is selected from the group consisting of acrylate- and methacrylate-functional
25 polymers.
5. A restorative system as in Claim 1 wherein the particulate additive is selected from the group consisting of metal salts of acrylates and methacrylates.
30
6. A restorative system as in Claim 1 wherein the particulate additive comprises nonmetallic filler comprising silane-treated pyrogenic silica.

- 20 -

7. A restorative system as in Claim 1 wherein the particulate additive comprises silanized zirconia/silica filler.

5 8. A restorative system as in Claim 1 wherein the particulate additive comprises an oxidizing agent.

9. A restorative system as in Claim 1 wherein the particulate additive comprises a reducing agent.

10

10. A restorative system as in Claim 4 wherein the particulate additive comprises poly(alkanoic acid) powder comprising acrylate- or methacrylate-functional groups.

15

11. A restorative system as in Claim 5 wherein the particulate additive is selected from the group consisting of zinc dimethacrylate, zirconium dimethacrylate, silver methacrylate, sodium methacrylate and magnesium methacrylate.

20

12. A restorative system as in Claim 8 wherein the particulate additive comprises benzoyl peroxide.

25 13. A restorative system as in Claim 9 wherein the particulate additive comprises sodium benzenesulfinate.

14. A restorative system as in Claim 2 wherein the modified amalgam alloy powder further comprises copper in about 2 to 30 percent by weight.

30

15. A restorative system as in Claim 14 wherein the modified amalgam alloy powder comprises "DISPERSALLOY" alloy powder and said particulate additive.

35

- 21 -

16. A restorative system as in Claim 3 wherein the dental adhesive is selected from the group consisting of "SCOTCHBOND" Dual Cure Dental Adhesive, "SCOTCHBOND 2" Light Cure Dental Adhesive, "PRISMA UNIVERSAL BOND 2" Dentin/Enamel Bonding Agent, "MIRAGE-BOND" Dentin-Enamel Bonding System, "RESTOBOND 3" Dual Dentin-Enamel Bonding Agent, "TENURE" Solution Dentin Bonding System, "GLUMA" Bonding System, "ALL-BOND" Universal Dental Adhesive System, "CLEARFIL" Photo Bond Light Cured Dental Bonding Agent, "BONDLITE" Dental Adhesive, "Johnson & Johnson" Dentin Bonding Agent, "Johnson & Johnson" Light Curing Bonding Agent, "PALFIQUE" Bonding Agent, "SHOFU" Bonding Base and "SINTERBOND" Dental Adhesive.

17. An amalgam composition of the type having an alloy powder amalgamated with mercury to form an amalgam restoration for carious tooth structure, wherein the improvement comprises:

particulate additive admixed with said alloy powder to form a modified amalgam upon amalgamation with mercury, wherein said particulate additive renders said modified amalgam adherent to tooth structure precoated with an acrylate- or methacrylate-functional dental adhesive.

18. An amalgam composition as in Claim 17 wherein the particulate additive is selected from the group consisting of acrylate- and methacrylate- functional polymers, metal salts of acrylates and methacrylates, nonmetallic fillers, oxidizing agents and reducing agents.

19. A restorative system as in Claim 1 wherein said particulate additive has the property of rendering said modified amalgam adherent to said tooth structure precoated with said dental adhesive such that the average

- 22 -

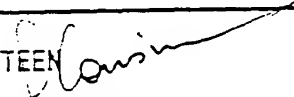
adhesive shear bond strength between the modified amalgam and precoated tooth structure is about 15 kg/cm² to 72 kg/cm².

- 5 20. An amalgam composition as in Claim 17 wherein said particulate additive has the property of rendering said modified amalgam adherent to tooth structure precoated with said dental adhesive such that the average adhesive shear bond strength between modified
10 amalgam and precoated tooth structure is about 15 kg/cm² to 72 kg/cm².

INTERNATIONAL SEARCH REPORT

PCT/US 91/08809

International Application No

I. CLASSIFICATION OF SUBJECT MATTER (If several classification symbols apply, indicate all) ⁶		
According to International Patent Classification (IPC) or to both National Classification and IPC		
Int.Cl. 5 A61K6/05; A61K6/00		
II. FIELDS SEARCHED		
Minimum Documentation Searched ⁷		
Classification System	Classification Symbols	
Int.Cl. 5	A61K	
Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in the Fields Searched ⁸		
III. DOCUMENTS CONSIDERED TO BE RELEVANT⁹		
Category ¹⁰	Citation of Document, ¹¹ with indication, where appropriate, of the relevant passages ¹²	Relevant to Claims No. ¹³
X	US,A,3 513 123 (J.SAFFIR) 19 May 1970 cited in the application see column 6, line 57 - line 72; claims; examples	1-2
Y	US,A,4 001 483 (H.LEE) 4 January 1977 cited in the application	1-3,6-7, 17-18
Y,P	EP,A,0 413 174 (MITSUBISHI RAYON CO.) 20 February 1991 see page 7, line 26	1-3,6-7, 17-18
Y	US,A,2 991 176 (J.CLANCY) 4 July 1961 cited in the application see column 1, line 18 - line 31	1-3,6-7, 17-18
A	US,A,4 064 629 (STONER) 27 December 1977 -/-	
<p>¹⁰ Special categories of cited documents:</p> <p>"A" document defining the general state of the art which is not considered to be of particular relevance</p> <p>"E" earlier document but published on or after the international filing date</p> <p>"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)</p> <p>"O" document referring to an oral disclosure, use, exhibition or other means</p> <p>"P" document published prior to the international filing date but later than the priority date claimed</p> <p>"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention</p> <p>"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step</p> <p>"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art</p> <p>"&" document member of the same patent family</p>		
IV. CERTIFICATION		
Date of the Actual Completion of the International Search	Date of Mailing of this International Search Report	
08 APRIL 1992	1 6. 04. 92	
International Searching Authority	Signature of Authorized Officer	
EUROPEAN PATENT OFFICE	G COUSINS-VAN STEEN 	

III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)

Category *	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claims No.
A	EP,A,0 291 944 (DEGUSSA AKTIENGESELLSCHAFT) 23 November 1988 & US,A,4 859 412 22 August 1989 cited in the application ---	

**ANNEX TO THE INTERNATIONAL SEARCH REPORT
ON INTERNATIONAL PATENT APPLICATION NO. US 9108809
SA 55181**

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on
The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information. 08/04/92

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
US-A-3513123	19-05-70	None	
US-A-4001483	04-01-77	None	
EP-A-0413174	20-02-91	JP-A- 3074310	28-03-91
		AU-B- 621233	05-03-92
		AU-A- 5795290	14-02-91
US-A-2991176		None	
US-A-4064629	27-12-77	AT-B- 353418	12-11-79
		AU-B- 504819	01-11-79
		AU-A- 2162277	03-08-78
		CA-A- 1090502	02-12-80
		DE-A- 2702923	11-08-77
		FR-A- 2338696	19-08-77
		GB-A- 1565441	23-04-80
		JP-A- 53004393	14-01-78
		NL-A- 7700678	28-07-77
		SE-A- 7700772	27-07-77
EP-A-0291944	23-11-88	DE-C- 3717048	03-11-88
		AU-B- 594574	08-03-90
		AU-A- 1652688	24-11-88
		DE-A- 3865087	31-10-91
		US-A- 4859412	22-08-89
US-A-4859412	22-08-89	DE-C- 3717048	03-11-88
		AU-B- 594574	08-03-90
		AU-A- 1652688	24-11-88
		DE-A- 3865087	31-10-91
		EP-A, B 0291944	23-11-88

